

Physical Properties of Nano-HAs/ZrO₂ Coating on Surface of Titanium Materials Used in Dental-implants and its Biological Compatibility

Pang Xiao-Feng*

Institute of Physical Electron and Life Science and Technology, University of Electronic Science and Technology of China, Chengdu 610054, China

*Corresponding author: Pang Xiao Feng, Professor, Institute of Life Science and Technology, University of Electronic Science and Technology of China, Chengdu 610054, China, Tel: (86) 28-83202595; E-mail: pangxf2006@yahoo.com.cn

Abstract

A gradient composite coating on the surface of titanium materials, which are used in dental implants, is prepared using an electric-chemical method. The physical properties of the composite coating and its strength of combining with titanium material are studied by the scanning electron microscope, energy dispersive spectrum and X-ray diffraction analysis, etc. The results show that the complex of nanohydroxyapatite/ZrO₂ is uniformly deposited and distributed on the surface of titanium materials; its strength of combining with titanium surface can reach 16.3 MPa, which is obtained using the tensile test. The immersion experiment shows that the new matter of carbonate-apatite is occurred and distributed uniformly on the surface of composite coating of nanohydroxyapatite/ZrO₂. The cell experiment of cultivate exhibits that the osteoblasts MG-63 can be grown well on the surface of the composite coating. These results indicate that the nanohydroxyapatite/ ZrO₂ composite coating on the surface of titanium materials has a good biological activity and compatibility, which could be used in the dental-implants.

Keywords: Nanohydroxyapatite; ZrO₂; Titanium materials; Osteoblasts MG-63; Electrochemical deposition method; Composite coating; Biological compatibility

Introduction

It is well known that the features of teeth in men and women can be changed with increasing age, in which some teeth could be damaged. Thus the dental-implants, which are implanted at the positions of damaged teeth to replace the latter, are developed and used widely in dentist medicine. At present, the dental-implants are, in general, manufactured by titanium materials. Clinic experiments show that the kind of dental-implants cannot be very well incorporated with the dental substrates, thus their biological functions will fail after about ten years. Therefore, it is very necessary to improve and lift the qualities of the dental-implants for enhancing the combined strength of the titanium materials with the dental substrates. The investigations show that nanomaterials, such as Hydroxyapatites (HAs), and their complexes with other nanomaterials, which are coated on the surface of titanium materials, can be used to enhance the incorporated strength with the dental substrates.

As it is known, the hydroxyapatite [Ca₁₀(PO₄)₆(OH)₂] (HA) is a mainly inorganic composition, which exists widely in bone and tooth in vertebrates and persons, and has an excellent biocompatibility. HAs can also incorporate with all biological tissues in virtue of interface reaction and hydrogen bonds^[1-7]. HAs can release the free calcium and phosphorus through the degradation, when are in a solution of organism, thus HAs can be easily absorbed by the biological tissues to stimulate the growth of tissues, bone and tooth in living system. Therefore, HAs have a wide applications as a repairing material in biological hard tissues.

Yang Yunzhi, *et al.* prepared the HAs coating on the surface of titanium alloy and measured its combined strength and toughness with the titanium alloy and inspected further its biological functions through injecting it into the tissues^[8]. Their results show that the combined strength of HAs with the biolog-

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ical tissues is increased relative to that of titanium alloy. Hence, they thought that HA coating could be utilized in the repairing of bone and tooth. However, the experimental measurements display that the combined strength of HA with the titanium alloy and its crystallization on the surface of the latter are lower^[9,10], then HAs are easily fallen off under action of a force or dissolved through the erosion effect in solution. Therefore the lifetime of use of the coating is very short. This phenomenon is due to the fact that the mechanical properties of HA, such as the thermal expansion coefficient, compressive strength, modulus of elasticity, flexural strength, hardness and fracture toughness, match not with those of the titanium materials. However, if some ceramic powders, such as nano-ZrO₂, are added into the coating to form a composite coating on the surface of titanium material, then the above shortcoming could be depressed or eliminated^[11].

Experimental measurements show that the thermal expansion coefficient and fracture toughness of ZrO₂ are just between those of HA and titanium material. At the same time, ZrO₂ has a good abrasion resistance, physiological corrosion - resistance and biocompatibility, its modulus of elasticity, flexural strength and hardness are also higher relative to those of HA and titanium alloys, which are shown in Table 1. Therefore, ZrO₂ can well combine with both the titanium materials and HA. If the HA/ZrO₂ composite coating on the surface of titanium alloys can be prepared, then the combined strength of the composite coating with the titanium material will be greatly enhanced relative to that of HA coating with the titanium material. Thus the combined strength of dental-implant with the dental substrates also will be greatly increased, when it is used in the dental-implants^[12-19]. Hence, it is quite necessary to study and prepare the HA/ZrO₂ composite coating on the surface of titanium materials for developing new biological ceramic materials used in bone and tooth.

Table 1: The mechanical properties of materials of HA, Ti₆Al₄V alloy and ZrO₂.

material	modulus of elasticity (GPa)	compressive strength (MPa)	flexural strength (MPa)	hardness (Hv/GPa)	fracture toughness (MPam ^{1/2})	expansion index (10 ⁻⁶ /K)
HA	81.4	462 ~ 509	103 ~ 113	3.43	0.69 ~ 0.96	11 ~ 14
Ti ₆ Al ₄ V	110 ~ 114	825 ~ 869	895 ~ 930	3.8 ~ 4.7	6 ~ 10	8.9
ZrO ₂	210	---	1000 ~ 1200	12.5 ~ 14.7	7 ~ 15	10.4

However, the experiments show that if ZrO₂ is only mixed into HA coating, instead of is injected between the HA and surface of titanium materials, then the biological effect of the composite coating is also not effective. Therefore, we should prepare the composite coating, which is composed of HA/ZrO₂, in which the ZrO₂ is injected between the HA and surface of titanium materials to form a new layer. In this case it is very necessary to improve the prepared method of the coating.

In this paper we use an electrochemical method to prepare the gradient composite coating of HA/ZrO₂ on the surface of titanium alloys through choosing appropriate electric voltage and current. This method has the following advantages.

(1) A gradient and dense composite coating of HA/ZrO₂ can be formed by this method, in which HA and ZrO₂ are orderly deposited on the surface of the titanium, i.e., ZrO₂ is uniformly between the HA and titanium.

(2) The temperature of the solution, in which the chemical reaction occurs, is lower, thus the phase transition and brittle cracking of the coating occurred in the prepared process can be avoided and eliminated by accommodating the electric voltage and current.

(3) The electric current and voltage of instrument, pH value of the solution, the thickness and uniformity of coating can be easily controlled in prepared process for a good and excellent bio-active coating.

(4) The materials, which are used in the chemical reaction, can be injected continuously, thus there are not pollutants in the experimental process. Therefore, it can eliminate the difficulties occurred in other methods.

Materials and Methods

Materials and Instruments

Materials used in this research, including LK2005A-type electrochemical workstation made by Nanlinke Co. CN, Scanning Electron Microscopy (SEM) with EDAX spectrum (Sanyo, JPN), X-ray diffractometer (GFL, Germany), 670-FT-IR spectrometer (Nicolet Nexus, USA) and electronic tensile tester (Shanghai, CN), are as follows: the calcium nitrate with four water molecules (Sigma, USA), ammonium dihydrogen phosphate (Sigma, USA), Hydrated nitric acid oxygen zirconium (Sigma, USA), Osteoblast cells (MG63) (Shanghai, CN), pure titanium material with purity of > 99.9% (Baoji British knight medical titanium Co. LTD, CN).

Experimental Method

The treatment of titanium pieces: The pure titanium materials are processed as some titanium slices of 1 cm × 1 cm × 0.1 cm with smooth surface, which are again washed and cleaned with ethanol, acetone, deionized water and ultrasonic wave about 15 mins, respectively. After this the titanium slices are further corroded in acid solution of HF:HNO₃:H₂O = 1:1:15 about 20 ~ 40 mins, we extract them from the solution to clean by the deionized water and ultrasonic wave for 8 mins.

Preparation of gradient and dense composite coating of HA/ZrO₂ on surface of the titanium: The processed titanium slices, platinum chips and saturation Hg₂Cl₂ are served as the worked electrode, pair electrode and reference electrode in the electrochemical equipment, respectively. The nitric acid zirconium oxygen is first dissolved as a deposited solution of zirconium with the zirconium concentration of 0.0125 mol/L by a dilute

nitric acid solution, the pH value of deposited solution should be regulated to 2.3 by adding ammonia water, thus the electrolyte deposited solution is prepared, which is used in the electrochemical workstation. When the strength of electric current in the worked electrode is adjusted to 11.1 mA and the temperature of the electrolyte deposited solution is controlled to 25°C, then Zr(OH)₄ can be well deposited on the surface of the titanium slices. The prepared titanium slices containing the Zr(OH)₄ coating are extracted from this solution after the deposited time of 400s and are dried further. Subsequently, we finish second electrochemical deposition of HAs on the surface of Zr(OH)₄ in the electrolyte solution at the temperature of 65°C to form HA/Zr(OH)₄ composite coatings on the surface of the titanium slices, where the strength of electric-current of the worked electrode is adjusted to 1.1 mA and the deposited time is controlled to 900s. In this process the electrolyte solution is prepared by the calcium nitrate with four water molecules, dihydrogen phosphate and distilled water, in which the pH value is adjusted to 4.1, the ratio of Ca vs. P is Ca : P = 1.67, the concentrations of Ca²⁺ and PO₄³⁻ ions are controlled to 0.021 mol/L and 0.0125 mol/L, respectively.

The prepared HA/Zr(OH)₄ composite coatings on the surface of the titanium slices are placed into the sintering furnace with vacuum condition to sinter at high temperature of

650°C after they are washed by the distilled water and dried further by far-infrared heating. Thus we can obtain the nano-HA/ZrO₂ composite coatings on the surface of titanium slices after the sintering times of 2h, which are required and can be used in the dental-implants.

Measurement of combined strength of the composite coating with the surface of titanium: We used the electronic-tensile tester to detect the combined strength of the nano-HA/ZrO₂ composite coating with the surface of titanium, which is an average of measured values obtained from three points, its unit is Pa.

The immersion experiment: The nano-HA/ZrO₂ composite coating on the surface of titanium material is soaked in a simulated solution of life (SSL) with 40 mL, which is placed in a closed and disinfectant centrifugal tube, the temperature and pH value of the solution are controlled to 36.5°C and 7.4, respectively. The components and their concentrations in the SSL are basically same with those in the Human plasma (HBP), which is shown in Table 2. The immersed composite-coatings extracted from SSL after 7 days and 14 days are washed and cleaned with de-ionized water and further dried, respectively, their properties are inspected by the bio-chemical method and SEM, respectively.

Table 2: The components and their concentrations in (SSL) and HBP(mM).

Ion	Na ⁺	K ⁺	Mg ²⁺	Ca ²⁺	Cl ⁻	HCO ₃ ⁻	HPO ₄ ²⁻	SO ₄ ²⁻
SSL	142.0	5.0	1.5	2.5	147.8	4.2	1.0	0.5
HBP	142.0	5.0	1.5	2.5	103.0	27	1.0	0.5

The observation of morphology of coating surface adhering the osteoblast MG63:

(A) The culture of cells. The MG63 cells are grown in a 5% CO₂ enriched incubator at 25°C in RPMI1640 media (Hyclone, American) supplemented with 5% fetal calf serum (FCS, Biological Industries, BaiAn, China). Using microscopic inspected-method we can verify that there are no contaminating cells in the third passage of cell culture.

(B) The experimental process. In all experiments, 4 × 10⁵ cells per well were seeded into 60 well micro-plates and allowed to grow continually under the conditions described above. The 100 μL/well foster liquids containing the fetal calf serum and insulin liquid of concentration of 1 mL/250 mL are added into each well. Subsequently, the titanium slices adhering the HA/ZrO_{2n} composite coating are placed into the wells. The whole micro-plates with titanium slices are again shifted into the CO₂ enriched incubator at 37°C to promote the proliferation and adherence of the MG63 cells on the surfaces of the composite coatings. After 24 hours the biological samples extracted from the enriched incubator are washed by the phosphate cushion fluid, dried by heating, fixed again by 2.5% glutaraldehyde about 24 hours and dehydrated further by the ethanol of 30 - 100% concentration about 20 min, then we can inspect the features of vaccination and adhesion of the MG63 cells on the surface of the composite coatings by SEM. Thus we eventually can determine the biological activity of the composite coatings from the measured results.

Results and Discussion

The influences of deposited time on the features of Zr(OH)₄ / HA coating

When the electric current of the worked electrode in LK2005A-type electrochemical workstation is adjusted to 11.1mA, the nano-HAs can be deposited on the surface of Zr(OH)₄, which was, first of all, adhered on the surface of titanium material, to form a HA/Zr(OH)₄ composite coating in the electrolyte solution of calcium phosphate. Figure 1a shows the outline of the HA/Zr(OH)₄ composite coating at the deposited time of 800s, which is obtained by SEM. Obviously, the features of composite coating depend on the relative weight of components, pH value, temperature of the electrolyte solution, deposited time and size of electric current of the worked electrode. Figures 1(b) and 1(c) show the outlines of the composite coating at the electric-current of 11.1 mA and 1.1 mA and the deposited times of 700s and 900s, respectively. From these results we see that the nano-HAs have not covered the whole surface of Zr(OH)₄, if the deposited time is too short and current is greater, which are shown in Figure 1(b), but if the deposited time is longer and current is smaller, then a good composite coating can be obtained as shown in Figure 1(c), where a dense needle structure containing some gaps are occurred. This kind of composite coating is helpful for the growth and adhesion of the osteoblast. Thus it could be used in the dental-implants^[20]. In Figure 2(a) we exhibit the outline of HA/Zr(OH)₄ composite coating on the surface of titanium material at 1.1mA and 850s. The compositions containing in the composite coating are shown in Figure 2(b),

which is obtained by Energy Dispersive Spectrometer (EDS). We see from this figure that there are really Ca, P, Zr, O elements in the composite coating. Thus we can affirm that the HA/Zr(OH)₄ composite coating is truly formed on the surface of titanium material in this condition.

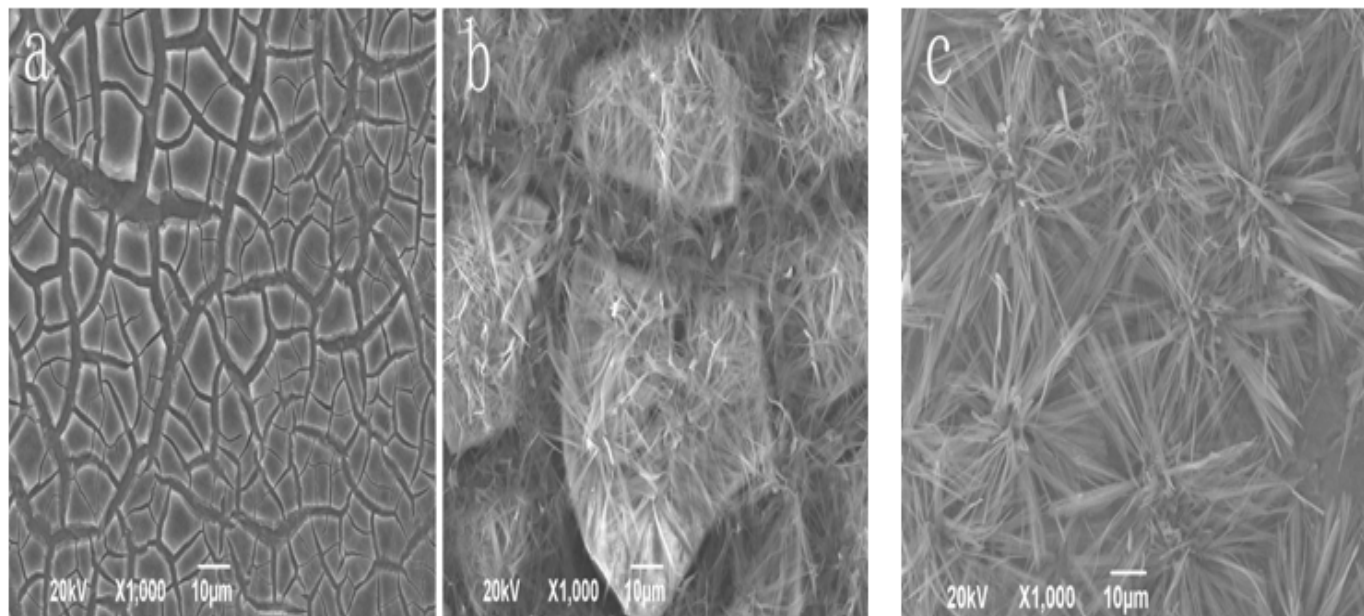


Figure 1: The outlines of HA/Zr(OH)₄ composite coating on the surface of titanium material at 11.1 mA, 800 s (a), 11.1 mA, 700 s (b) and 1.1 mA, 900 s (c), respectively.

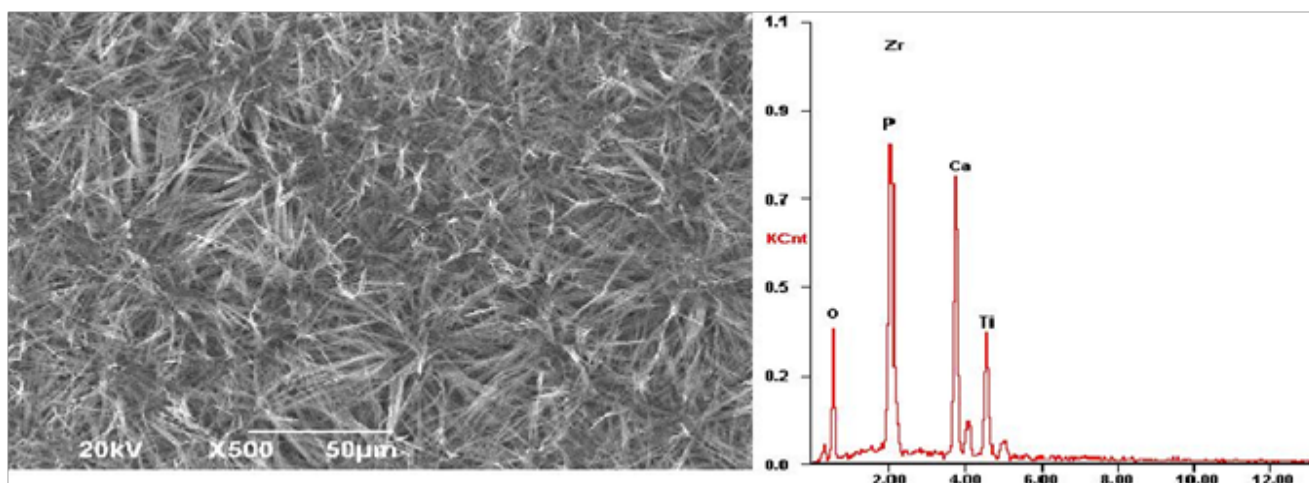


Figure 2: The outline of HA/Zr(OH)₄ composite coating on the surface of titanium material formed in the conditions of 1.1 mA and 850 s and its compositions obtained by EDS.

The influence of deposited temperature on the features of HA/Zr(OH)₄ composited coating

Experiments display that the features of HA/Zr(OH)₄ composite coating on the surface of titanium materials depend closely on the temperature of the electrolyte solution. We collect the changes of feature of HA/Zr(OH)₄ composite coating with increasing the temperature by SEM, which are shown in Figure 3. Figure 3(a) exhibits that the nano-HAs cover not the whole surface of Zr(OH)₄, which is early adhered on the surface of titanium materials, where the nano-HAs are non-uniformly gathered together to form some groups on the surface of the Zr(OH)₄ coating at lower temperature of 25°C. However, when the deposited temperature is lifted to 45°C, although the nano-HAs are covered the whole surface of Zr(OH)₄ coating as shown in Figure 3(b), its density is too great to depress its bio-activity. Figure 3(d) shows that the needle structure of nano-HAs is covered the whole surface of Zr(OH)₄ coating, but the clusters of nano-HAs having great area occurs also in the surface at the temperature of 80°C. However, an uniform and dense HA/Zr(OH)₄ composite coating having the needle structure on the surface of titanium materials is obtained at the temperature 65°C, which is shown in Figure 3(c). This structure is very advantageous for the osteoblast's growth and adhesion on its surface.

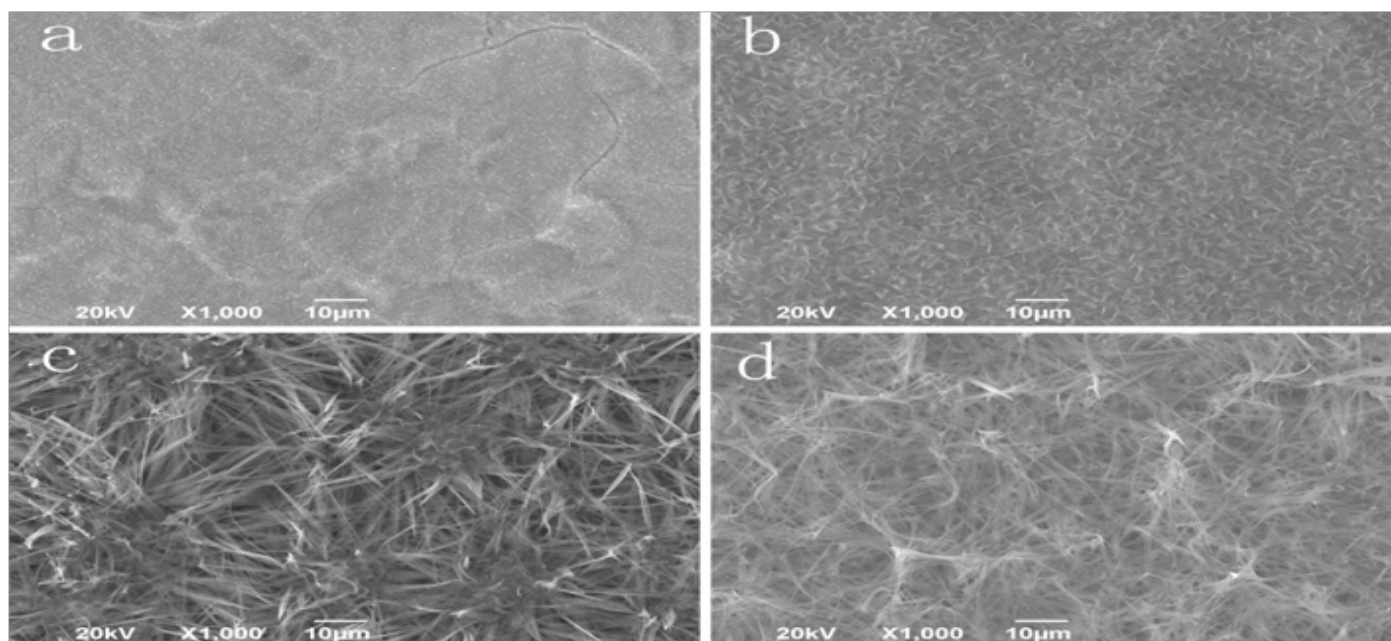


Figure 3: The SEM images of HA/Zr(OH)₄ composite coating at different temperatures of 25°C (a), 45°C (b), 65°C (c) and 80°C (d), respectively.

The changes of combined strength of HA/ZrO₂ composite coating with the surface of titanium material after sintering

Experiments show that HA/Zr(OH)₄ composite coatings can become HA/ZrO₂ composite coating and its combined strength with the surface of titanium material is increased after they are thermally sintered. We used the electronic tensile tester to detect the combined strength of HA/ZrO₂ composite coating with the surface of titanium material after sintering at 650°C, which is 16.3MPa, but the combined strength of nano-HA coating with the surface of titanium material after sintering at 650°C is only 5.8MPa. The values are an average of measured values obtained at three points as shown in Table 3. This shows clearly that ZrO₂, which is served as a transition layer, can well ease and offset the non-matched effect of thermal expansion coefficient of HA with that of the titanium, thus the combined strength of HA/ZrO₂ composite coating is significantly enhanced due to the match of thermal expansion coefficient between them. In Figure 4 we exhibit the XRD pattern of HA/ZrO₂ composite coating on the surface of titanium material after thermal sintering at 650°C, which shows clearly that the nano-HAs are already covered on the surface of ZrO₂ coating. In Figure 4 there are two strong peaks, which are due to the large thickness of the composite coating about 3 μm obtained by the ladder meter, and some weaker peaks, which denote the existence of precursors CaHPO₄ of HAs in the coating arising from the lower of sintered temperature, in the titanium substrates.

Table 3: The combined strength of HA and ZrO₂/HA coatings after thermal sintering.

Sample	HA coating	HA/ZrO ₂ coating (after sintering)
1	4.7	14.6
2	5.6	16.1
3	7.1	18.3
average	5.8	16.3

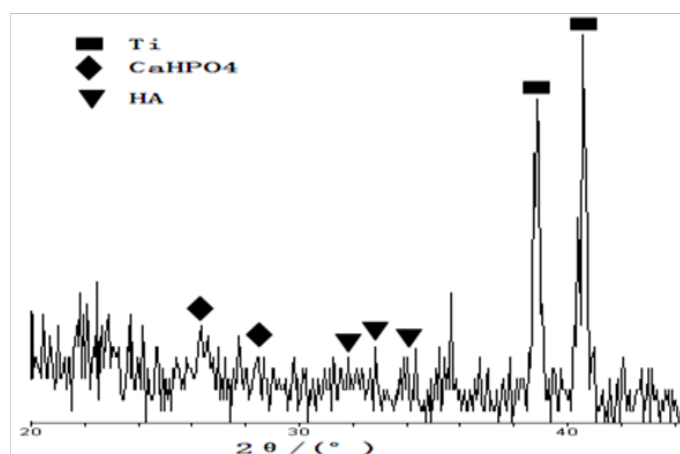


Figure 4: XRD pattern of composite coating after sintering.

The biological activity of the composite coating

We use the immersion method and SEM to study and inspect the biological activity of the nano-HA/ZrO₂ composite coating on the surface of titanium. Figures 5a and 5b are the outlines of Zr(OH)₄ coating and nano-HA/ZrO₂ gradient coating before the immersion, respectively, figures 5c and 5e are the features of Zr(OH)₄ coating at different immersion times of 7 days and 14 days, respectively, which show that NaCl grains are formed and accumulated on the surface of the Zr(OH)₄ coating, and the accumulation of NaCl is also increased with increasing the immersion time, in which but there are not growth of apatite particles induced by SSL, thus the apatite particles are not covered on the surface of the coating. Figures 5d, 5f and 5g exhibit the outlines of nano-HA/ZrO₂ gradient coating at different immersion time of 7 days and 14 days, respectively. Figure 5d shows that there are some clusters formed by the apatite particles of white ball in the composite coating, the size of apatite particle is about 10 μm, they cover completely over the whole surface of ZrO₂ coating. However, there are also some small cracks in the coating due to contraction of the surface arising from the drying of the coating after they are extracted from the immersion solution. Figure 5f indicates that the sizes of apatite particles and

thickness of coating are increased in the coating in this condition. In the meanwhile, the blocky structure composed of small apatite balls is also appeared in the coating, when the immersion time is increased, which is 14 days. Figure 5g is a magnified result of pattern of HA/ZrO₂ gradient coating after immersion time of 14 days, which is obtained by SEM with high resolution. These results indicates clearly that plenty of small needle HAs can be grown from the surface of a great spherical-bodies, which is accumulated by a lot of nano-HAs. Thus the mineralization and growth of apatite particles induced by SSL occur. These results manifest that the nano-HA/ZrO₂ gradient coating has a good biological activity.

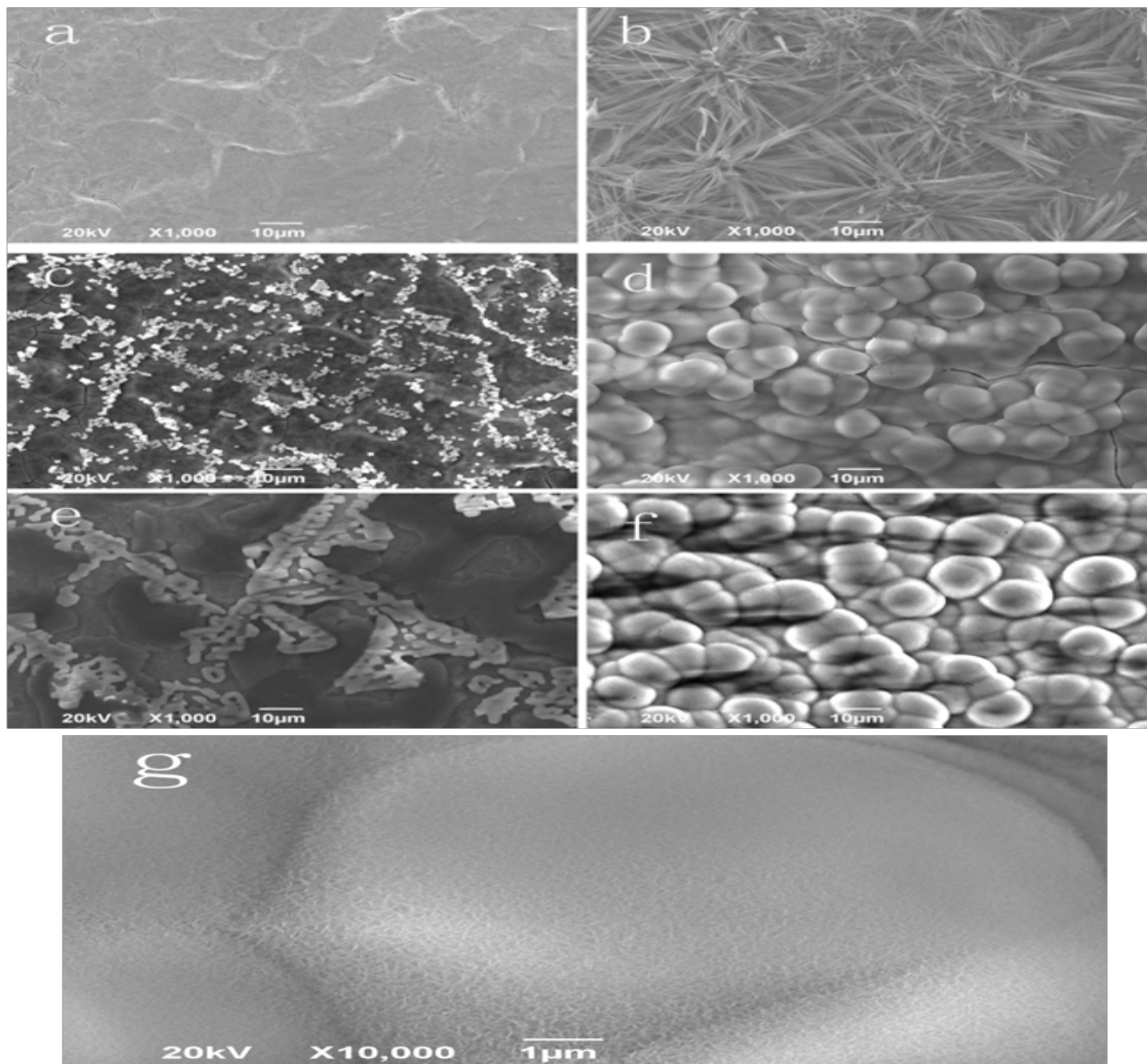


Figure 5: SEM images of Zr(OH)₄ coating and gradient coating after different immersion times.

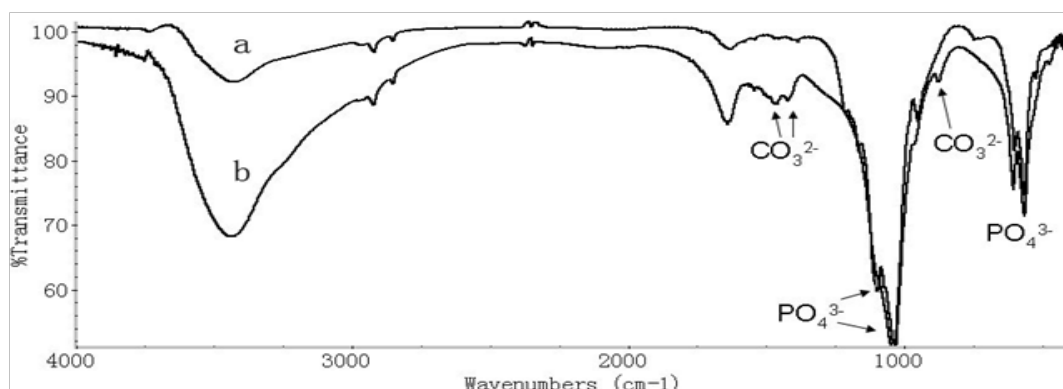


Figure 6: Infrared spectrum of absorption of nano-HA/ZrO₂ coating after the immersion time of 7 days in SSL.

We collected also the infrared spectrum of absorption of nano-HA/ZrO₂ composite coating on the surface of titanium after the immersion time of 7 days in SSL using 670-FT-IR spectrometer, which is shown in Figure 6. We see from this figure that some new peaks at 1489.38, 1412.92, 871.59 cm⁻¹ occur after the immersion, which correspond to the characteristic peaks of CO₃. This means that the new matter of carbonate-apatites, such as like-bone apatites, which are formed in the composite coating arising from SSL in the immersion process, which induces the mineralization and growth of bone apatites^[21,22]. This demonstrates again that nano-HA/ZrO₂ gradient coating is existed in this case.

The biological compatibility of nano-HA/ZrO₂ composite coating

We further study the biological compatibility of nano-HA/ZrO₂ composite coating in virtue of inspecting of growth states of osteoblast MG63 on its surface. In this experiment the nano-HA/ZrO₂ coatings on the titanium pieces are together placed in the culture solution of MG63 cells. Figure 7 shows that the states of proliferation and adherence of osteoblasts MG63 on the surfaces of the pure titanium and nano-HA/ZrO₂ coatings after culture of 24h, respectively. Figure 7b shows that MG63 cells are adhered well on the surface of nano-HA/ZrO₂ coatings, the

MG63 cells are proliferated along these needle structures, which provide more adherent positions for the proliferations of cells. Meanwhile, we see that the adhered areas of cells are gradually extended to form a structure of anchor shape in these gaps in the composite coating. This indicates that the nano-HA/ZrO₂ composite coating has a good biological compatibility. However, the MG63 cells cannot proliferate on the surface of pure titanium pieces as shown in Figure 7a. This verified again that the dental-implants made by the pure titanium pieces have not a good biological compatibility as mentioned in introduction.

Modern cell biology and medicine indicate that when the dental-implants are implanted on the substrates of teeth, the plasma proteins will be adhered on surface of the dental-implants to form a molecular layer, which can absorb the osteoblasts and can extend their proliferation on the surface of dental-implant in virtue of the plasma proteins. Thus the dental-implants are fixed on the substrates of teeth. This represent clearly that the dental-implants made by the titanium adhering the nano-HA/ZrO₂ composite coating are easily and firmly fixed on the substrates of teeth because the composite coating can provide more adherent positions for the cells through these needle structure containing some gaps. Therefore, the titanium adhering the nano-HA/ZrO₂ composite coating prepared by us have a good biological compatibility, thus a wide application in dental medicine.

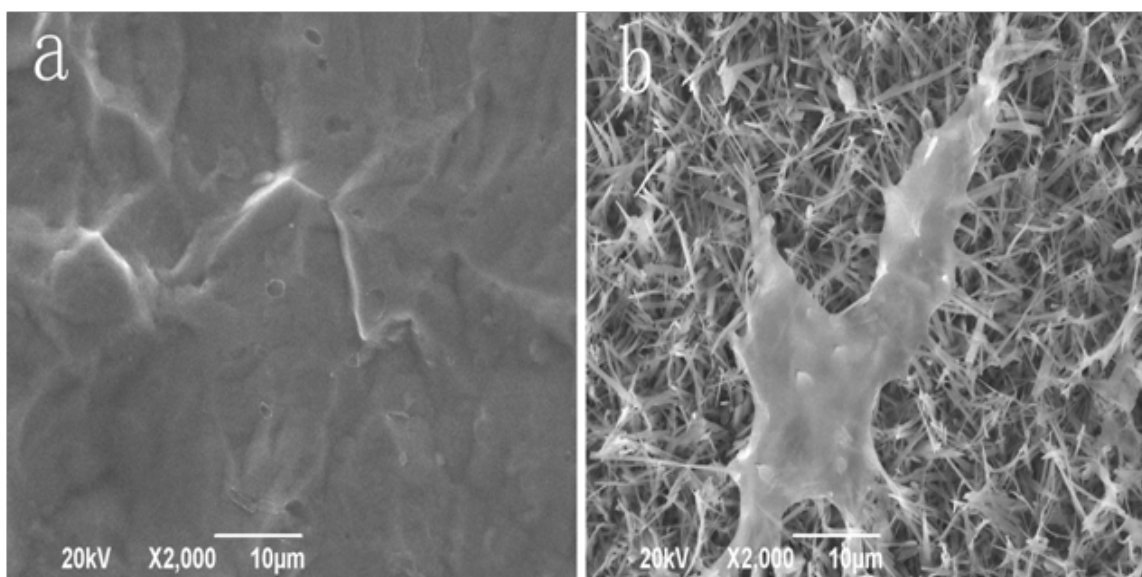


Figure 7: SEM images of proliferation of MG 63 cells on the surfaces of pure titanium coating (a) and composite coating (b), respectively

Conclusion

The HA/ZrO₂ composite coating on the surface of titanium materials, which are used in dental implants in dentist medicine, are prepared by the electrolyte solution composed of ZrO(NO₃)₂, Ca(NO₃)₂ and NH₄H₂PO₄ using an electric-chemical method, the physical properties including the combined strength and biological activity and compatibility of the composite coatings are studied by methods of SEM, EDS spectrum, X-ray diffraction and infrared spectrum of absorption, etc.. From these investigations we can draw the following conclusions.

(1) The nano-HA/ZrO₂ composites can be uniformly deposited and adhered on the surface of titanium materials, its combined strength with the surface of titanium is very high, and can reach 16.3 MPa.

(2) The immersion experiment in SSL shows that the carbonate-apatites can be uniformly deposited on the surface of HA/ZrO₂ composite coating. This means that the composite coating has a good biological activity.

(3) The cell cultivate experiment denotes that the MG63 cells can be adhered well on the surface of nano-HA/ZrO₂ coatings, where the MG63 cells can also proliferate along these needle structures, which provide more adherent positions for the proliferation of cells. Meanwhile, the adhered areas of cells are gradually extended to form a structure of anchor shape in these gaps in the composite coating. This indicates that the nano-HA/ZrO₂ composite coating has a good biological compatibility. Thus, we can predict that the titanium adhering the nano-HA/ZrO₂ composite coating prepared by us have a wide application in dental medicine.

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